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- (56) References cited: EP-A- 0 286 903

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- ARCH . OPHTHALMOL. vol. 108, no. 8, 1990. pages 1102 - 1105 J. VILLUMSEN 'The effect of adding prostaglandin F2alpha-isopropylester to
- timolol in patients with open angle glaucoma." · OPHTHALMOLOGY vol. 98, no. 7, 1991, pages 1079 - 1082 P.-Y. LEE 'Additivity of prostagiandin F2alpha-1-isopropyl ester to timolol in glaucoma patients."

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Description

BACKGROUND OF THE INVENTION

1. Field of the Invention

[0001] The present invention relates to the treatment of ocular hypertension with atternate administration of (a) a βadrenergic blocker and (b) a prostanoic acid compound with an improved efficiency.

[0002] The compounds used as the component (b) in the present invention are prostaglandin analogues.

2. Information of Prior Art

[0003] It is well known that the production and effluence of the aqueous humour, which are the important factors for the circulation of the aqueous humour, and hence the intraocular pressure as the results thereoft, vary with the circadian rhythm. Generally, in humans, the phase in which the production enhances is the daytime, during which the production of the aqueous humour is faithlisted and the intraocular pressure rises. On the other hand, the phase in which the aqueous humour is childred and the intraocular pressure fails. In contrast, in reables, the phase in which the aqueous humour production enhances is the night and the phase in which the aqueous humour production enhances is the night and the phase in which the aqueous humour production enhances is the night and the phase in which the aqueous humour production supresses is the daytime, 10004. This circadian rhythm of the intraocular pressure is observed not only in healthy humans but also in subjects of coular hypertension such as with glaucoma and a possibility that a relatively big variation in the intraocular pressure is a continuous need for the development of an improved method for treatment of ocular hypertension in which the intraocular pressure is effectively contributed laking the circadian rhythm of ocular tension in the hypertensive subjects.

into consideration.

(2005) The β-adrenergic blockers are the most widely used drugs for the treatment of glaucoma and ocular hypertension. In a report studying a relation between the circadian thythm of intraocular pressure and the ocular hypotensive activity of Timolius as significant in the enhancement phase of aqueous humour production, i.e. daytime in humans and night in rabbits, but negligible in the suppression phase of aqueous humour production, le. night in humans and dispit in arbbits, but negligible in the suppression base of particular than the parent (or observable) effect of β-adrenergic blockers such as Timolo is high at the enhancement phase of aqueous humour production and low at the suppression phase of aqueous humour production and low at the suppression phase of aqueous humour production and low at the suppression phase of aqueous humour production are acqueous humour and that controlling of the ocular tension is important for the treatment of ocular hyperension also in the suppression phase of aqueous humour production, it is considered that treating the ocular hyperension with a β-adrenergic blocker only is insufficient.

[0006] Prostanoic acid refers to the basic skeleton, shown by the formula below, as the common structural feature of the naturally occurring prostaglandins (hereinafter, prostaglandins are referred to as PGs).

The primary PGs are classified based on the structural feature of the five-membered cycle molety into PGAs, PGBs, PGCs, PGDs, PGBs, PGGs, PGBs, PGHs, PGBs and PGJs, and also on the presence or absence of unsaturation and oxidation in the chain molety as

55 Subscript 1 - - - 13,14-unsaturated-15-OH

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Subscript 2 - - - 5.6- and 13,14-diunsaturated-15-OH

Subscript 3 - - - 5,6- 13,14- and 17,18-triunsaturated-15-OH

Further, PGFs are sub-classified according to the configuration of hydroxy group at position 9 into α (hydroxy group being in the alpha configuration) and β (hydroxy group being in the beta configuration).

[0007] The fact that the above compounds under item (b) have ocular hypotensive activity has been known by Japanese Patent Publication No. A-108/1980. It has also been cescribed in Japanese Patent Publication No. A-313728/1988, page 7, column 3, lina 7 rom bottom to page 8, column 4, line 4, that a combination of PGF₂ is ejectroyly aster and Timolo (an agent for treating glaucoma) may be advantaged because the ocular hypotensive activity of the former is not inhibited by a β-adrenergic blocker such as the latter, Furthermore, a synergistic combination of a β-adrenergic blocker and a 13,4-dhypfor-15-sten OFE is described in EP-A-46569 (thev. 27, 1991). Such description, however, does not suggest that an alternate use of the β-adrenergic blocker and the component (b) in the present invention gives improved results.

[0008] Similarly, a synergistic combination of a β -actrenergic blocker and a POF_{2n} ester is described in EP-A-0 286 903. Ophthalmology 80 (2), 1916, 1079; and Arch. Ophthalmology 10 (8), 1990, 1102. Although a synergistic relationship between the two agents is shown, the documents of not suggest that alternate use of the two agents gives improved results.

5 [0009] After an extensive study the present inventor has surprisingly discovered that the prostancic acid compounds exhibit a significant ocular hypotensive activity at the surpression phase of aqueous humor production in which the β-adrenergic blockers such as Timolol can hardly exhibit the ocular hypotensive activity.

SUMMARY OF THE INVENTION

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[0010] In a first aspect, the present invention provides the use of a β -adrenergic blocker and of a derivative of prostancia cald for the manufacture of a therapeutic kill for the concomitant treatment of ocular hypertension, wherein the β -adrenergic blocker is to be administered only in the enhancement phase of aqueous humour production and the derivative of prostancia cald is to be administered only in the suppression phase of aqueous humour production

[0011] In a second aspect the present invention provides the use of a β-adrenergic blocker and of a derivative of prostancic acid for the manufacture of a therapeutic kit for the concomitant treatment of glaucoma, wherein the β-adrenergic blocker is to be administered only in the enhancement phase of aqueous humour production and the derivative of prostancic acid is to be administered only in the suppression phase of aqueous humour, production.

DETAILED DESCRIPTION OF THE INVENTION

[0012] The β-adrenergic blockers used in the present invention refer to agents capable of blocking the β-adrenergic receptor. Typical examples of such agents are relatively less selective β-adrenergic receptor blocking agents which are represented by the following formula:

A-OCH_CH(OH)CH_NHC(CH_)(R)

wherein A is an aromatic group and R is hydrogen atom or methyl.

[0013] The above group A includes 4-morpholino-1,2,5-thiadiazol-3-yl, 2-acetylbenzefuran-7-yl, 1,2,3,4-tetrahydro-2-oxo-quincline-5-yl. Preterred compounds include Timolol, Betundol, Betaxolol, Levabunciol, Carteciol and pharma-ceutically acceptable salls thered such as inorganic salls, e.g. hydrochloride or organic salls, e.g. maleete.
[0014] The term prostanoic acid compound refers to a compound (or derivative) in which one or more atom or group

(or molety) in the prostanoic acid shown by the formula (A) is replaced by other atom or group relimitated. Such is derivalization includes the modifications known in the synthetic PG analogues such as those shown below and other modifications. The prefetred prostanoic acid compounds have the ocular hypotensive activity and particularly aqueous humor effluence enhancing activity.

Nomenclature

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[0015] Nomenclature of the prostanoic acid compounds herein uses the numbering system of prostanoic acid represented in formula (A) shown above.

[0016] While formula (A) shows a basic skeleton having twenty carbon atoms, the compounds used in the present invention are not limited to those having the same number of carbon atoms. The carbon atoms in Formula (A) are numbered 2 to 7 on the c-chain starting from the c-action atom adjacent to the carbonytic carbon atom which is numbered 1 and towards the flive-membered ring, 8 to 12 on the said ring starting from the carbon atom adjacent to the ring. When the number of carbon atoms adjacent to the ring. When the number of carbon atoms is decreased in the c-chain, the number is deleted in order starting from position 2 and when the

number of carbon atoms is increased in the c-chain, compounds are named as substituted derivatives having respective substituents at position 1 in place of carbony group (C-1). Similarly, when the number of carbon atoms is decreased in the a-chain, the number is deteled in order starting from position 20 and when the number of carbon atoms is increased in the u-chain, compounds are named as substituted derivatives having respective substituents at position 20. Streechemistry of the compounds is the same as a that of above formula (A) unless otherwise specific

[0017] The above formula expresses a specific configuration which is the most typical one, and in this specification compounds having such a configuration are expressed without any specific reference to it.

[0018] In general, PGDs, PGEs and PGFs have a hydroxy group on the carbon atom at position 9 and/or 11 but the compounds used in the present invention includes PGs having a group other than a hydroxy group at position 9 and/or or 11. Such PGs are referred to as 9-dehydroxy-9-substituted-PG compounds or 11-dehydroxy-II-substituted-PG compounds

IDD19]. As stated above, nomenclature of the prostancia acid compounds in based upon the prostancia paid. These compounds, however, can also be named seconding to the IUPAD reming paytem. For swample, 13.14-d-Hybro-15-kato-1881,S-Huono- PGE, is: Q17-1(II-R,R,R,R)3-d-hydroy-2-(HR,S,Huono-3-oxo-1-oxyl)5-boxcepolopentyl)-hept.5-enote, cla. 13,14-d-Hybro-15-kato-19-mbyl-2-(HR,R,R,R)3-d-hydroy-2-(HR,R,R)3-d-hydroy-2-(HR,R)3-d-hydroy-1-(HR,R)3-Hydroy

Preferred Compounds

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(3-oxo-1-nonvi)-cyclopentyl)-hept-5-enonate.

[0020] Preferred prostanoic acid derivatives used in the present invention are those having an oxo group at position 15 of the prostanoic acid in plece of the hydroxy group as a feature. These derivatives may have as inglie bond (15-keto-PG, compounds), at double bond (15-keto-PG, compounds) between positions 5 and 6, or two double bonds (15-keto-PG, compounds) between positions 5 and 6 as well as positions of 17 and 18.

[0021] Examples of substitution products or derivatives include pharmaceutically or physiologically acceptable salts and esters at the carboxy group at the alpha chain, unsaturated derivatives having a double bond or a triple bond between positions 2 and 3 or positions 5 and 6, respectively, substituted derivatives having substituent(s) on carbon atom(s) at position 3, 6, 16, 17, 19 and/or 20 and compounds having lower allyl or a hydroxy (lower) allyl group at position 9 and/or 11 in place of the hydroxy group, of the above PGs.

[0022] Examples of substituents present in preferred compounds are as follows: Substituents on the carbon atom at position 3, 17 and/or 19 include lower alkyl, or example, C_{1,4} alkyl, especially methyl and ethyl. Substituents on the carbon atom at position 16 include lower alkyl e.g. methyl, ethyl etc., hydroxy and halogen atom e.g. chlorine, fluorine, aryloxy e.g. trill urromethyly hency, etc. Substituents on the carbon atom at position 20 include saturated and unsaturated lower alkyl e.g. C_{1,4} alkyl, lower alkoxy e.g. C_{1,4} alkyl, solven atoxy e.g. C_{1,4} alkyl, over alkyl or one of the carbon atom at position 6 include ox or group forming carbony. Stereochemistry of PGa having hydroxy, lower alkyl or lower (hydroxy) alkyl substituent on the carbon atom at position 9 and/or 11 may be alpha, beta or mixtures thereof.

[0023] Sald derivatives may have an alkoxy, phenoxy or phenyl group at the end of the omega chain where the chain is shorter than the primary PGs.

[0024] Especially preferred compounds are those having a lower alkyl such as methyl, ethyl, etc. at position 20.

[0025] A group of preferred compounds used in the present invention has the formula

$$\begin{array}{c}
X \\
R_1 - A \\
B - C - R_2 \\
Y \\
Z
\end{array}$$
(1)

wherein X and Y are hydrogen, hydroxy, halo, lower alkyl, hydroxy(lower)alkyl, or oxo, with the proviso that at least

one of X and Y is a group other than hydrogen, and 5-membered ring may have at least one double bond, A is -COOH or its pharmaceutically acceptable salt or ester, B is -CH₂-CH₂-, -CH=CH- or -C=C-, Z is

wherein R₃ is lower alkyl or acyl,

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PA, is bivalent saturated or unsaturated, lower or medium aliphatic hydrocarbon residue which is unsubstituted or substituted with halo, oxo or anyl.

R₂ is saturated or unsaturated, medium aliphatic hydrocarbon residue which is unsubstituted or substituted with halo, hydroxy, oxo, lower alkoxy, lower alkanoyloxy, cyclo(lower)alkyl, anyl or anyloxy.

- (9025) In the above formula, the term "unsaturated" in the definitions for Fi, and R₂ is intended to include at least one and optionally more than one double bond ander triple bond isolatedly, sepretally or serially present between carbon atoms of the main and/or alide chains. According to usual nonmerclainer, an unsaturation between two statis positions is represented by denoting the lower number of said two positions, and an unsaturation between two distal positions is represented by denoting the lower number of said two positions, and an unsaturation between two distal positions is represented by denoting both of the positions. Preferred unsaturation is a double bond at position 2 and a double or triple bond at position 5.
- 90 [0027] The ferm "lower or medium alighalic hydrocarbon residue" or "medium alighalic hydrocarbon residue" refers to a straight or branched chain hydrocarby topun having 1 to 1 4 carbon atoms re 5 to 14 curbon atoms, respectively, (for a side chain, 1 to 3 carbon atoms being preferred) and preferably 2 to 8 carbon atoms for R, and 6 to 9 carbon atoms.

[0028] The term *halo* denotes fluoro, chloro, bromo and iodo.

- [0029] The term "lower" throughout the specification is intended to include a group having 1 to 6 carbon atoms unless otherwise specified.
 - [0030] The term 'lower alkyl' as a group or a moiety in hydroxy(lower)alkyl, monocyclic ary(lower) alkyl, monocyclic aroy(lower) alkyl and beligner alkyl includes saturated and straight or branched chain hydrocarbon radicals containing 1 to 6, cerbon atoms, e.g. methyl, ethyl, propyl, sopropyl, butyl, sobutyl, butyl, propyl and hasyl.
- [0031] The term "lower alkoxy" refers to the group lower-alkyl-O- wherein lower alkyl is as defined above.
 - [0032] The term "hydroxy(lower)alkyl" refers to lower alkyl as defined above which is substituted with at least one hydroxy group, e.g. hydroxymethyl, 1-hydroxyethyl, 2-hydroxyethyl and 1-methyl-1-hydroxyethyl.
 - [0033] The term "lower alkanoyloxy" refers to a group of the formula: RCO-O- wherein RCO- is an acyl group formed by oxidation of a lower alkyl group as defined above, e.g. acetyl.
- 35 [0034] The term "cyclo(lower)alkyl" refers to a cyclic group formed by cyclization of a lower alkyl group as defined above.
 - [0035] The term "anyl" includes unsubstituted or substituted aromatic carbocyclic on heterocyclic (preferably monocyclic) groups, e.g.-phonyl, tolyl, xylyl and thilenyl. Examples of substituents are halo and halo(lower)alkyl wherein halo and lower alkyl beling as defined above.
- 40 [0036] The term "aryloxy" refers to a group of the formula: ArO- wherein Ar is anyl as defined above.
 - [0037] Suitable 'pharmaceurically acceptable salls' includes conventional non-toxic salts, and may be a salt with an inorganic base, for example an alkali metal salt (e.g. sodium salt, potassium salt, etc.) and an alkaline earth metal salt (e.g. calcium salt, magnesium salt, etc.) ammonium salt, as all with an organic base, for example, an amhe salt (e.g. calcium) salt, as all with an organic base, for example, an amhe salt (e.g. methylamine salt, dimethylamine salt, cycloherylamine salt, propridine salt, propridine salt, expenditum) salt, propridine salt, prop
- 45 ethanoismine salt, diethanoismine salt, triethanoismine salt, trist(hydroymethylammoyethne salt, monomethylamonethanoismine salt, procine salt, decl, a basic amino and salt (le g, arginine salt, eyeine salt, etc.), a basic amino and salt (le g, arginine salt, syeine salt, etc.) tetrasityl ammonium salt and the like. These salts can be prepared by the conventional process, for exemple from the corresponding acid and base or by salt interchange.
- [0038] Examples of the "pharmaceutically acceptable setters" are alliphatic extens, for example, lower alkyl exter e. 9, methyl exister, they extern proper sters, tonyor exters, tonyor extens, busyl exister, busylar exter, extensive propylethyl exter, exc., lower alkeryl exter, exc., lower alkeryl exter, exc., lower alkeryl exter, exc., and extensive except extensive extens
 - [0039] Preferred examples of A include -COOH, -COOCH₂, -COOCH₂CH₃ and -COOCH(CH₃)₃.
 - [0040] The configuration of the ring and the α- and/or omega chain in the above formula (I) may be the same as or

different from that in the primary PGs. However, the present invention also includes a mixture of a compound having a primary configuration and that of an unprimary configuration. [0041] A group of more preferred compounds used in the present invention has the formula

$$\begin{array}{c}
X \\
R_1 - A \\
B - C O - R_2
\end{array}$$
(II)

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wherein X and Y are hydrogen, hydroxy, halo, lower alkyl, hydroxy(lower)alkyl, or oxo, with the proviso that at least one of X and Y is a group other than hydrogen, and 5-membered ring may have at least one double bond, A is <COOH or its pharmaceutically acceptable sail or ester, B is <CH₂-CH₂-CH-CH- or <C=C-C. B, is bitvaient saturated or unsaturated, lower or medium aliphatic hydrocarbon residue which is unsubstituted or substituted with halo, tox or anyl, B; is saturated or unsaturated, medium aliphatic hydrocarbon residue having 5 or more carbon stores in the main or straight chain molety which is unsubstituted or substituted with halo, hydroxy, oxo, lower alkoxy, lower alkancyloxy, cyclo(lower)alkyl, anyl or anyloxy.

[0442] Examples of the typical compounds of the present invention are 15-keto-20-lowerally-PGA-Fs and their a2derivalives, SR, Smethyl-derivalives, 6-rox-derivalives, SRS-fution-orderivalives, 1618-fution-orderivalives, 1618-fution-ord

[0043] The compounds having 15-keto group may be in the keto-hemiacetal equilibrium by forming a hemiacetal between hydroxy group at position 11 and ketone at position 15.

[0044] The proportion of both tautomoric isomers, when present, varies depending on the structure of the rest of the molecule or kind of any substitute present and, sometimes, one isomer may precioninally be present in comparison with the other. However, in this invention, it is to be appreciated that the compounds used in the invention include both isomers. Further, while the compounds used in the invention include both isomers. Further, while the compounds used may be represented by a structure or name based on kato form reparties of the presence or absence of the isomers, it is to be noted that such structure or name does not intend elimination of the hermiscella layer of compounds.

[0045] In the present invention, any of the individual tautomeric isomers, a mixture thereof, or optical isomers, a maxture thereof, a racemic mixture, and other isomers such as steric isomers can be used in the same purpose. [0046] Some of the compounds used in the present invention may be prepared by the method disclosed in Japaneses

Patent Publications (unexamined) No. A-108/1990 and A-96528/1990.

(0047) Alternatively, these compounds may be prepared by a process analogous to that described in the above publications in combination with the known synthetic method for the five-membered ring moiety.
(0048) In the process for preparing 13,14-dillydro-15-keto-compound:

A commercially available (-)-Corey (actone, which is used as a starting material, is subjected to Collins oxidation to give an aidehybe. The aidehyde is allowed to reach with dimethyl (2-coadisylphosphomate anion to give an a, g-un-saturated ketone, and the resultant is reduced to ketone. The carbonyl group of the ketone is allowed to react with a colling to the commercial control of the control of the commercial control of the commercial control of the commercial control of the commercial control of the control o

[0049] Using the above tetrapyranyl ether as a starting material, 6-keto-PG₁s of the formula:

may be obtained as follows:

The letrapyranyl ether is reduced using disobutyl aluminium hydride and the like to give a lactol, which is allowed to react with a yilde obtained from (4-carboxybutyl)tripnerylphosphonium bromide, and the resultant is subjected to estertication followed by cyclization, combining he 5.6-double bond and the C-9 hydroxyl group with NBS or lodine, providing a halide. The resultant is subjected to dehydrohalogonation with DBU and the like to give a 6-keto compound, which is subjected to Jones oxidation followed by deprotection to give the objective compound.

may be obtained as follows:

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The above tetrapyranyl ether is reduced to the lactol, which is allowed to react with a yilde obtained from (4-carboxybutyl)tiphenylphosphonium bromide to give a carboxylic acid. The resultant is subjected to esterification followed by Jones oxidation and deprotection to give the objective compound. [0051] In order to obtain PG,s of the formula:

using the above tetrapyranyl ether as a starting material, in the same manner as PG2 of the formula:

the 5,6-double bond of the resulting compound is subjected to catalytic reduction followed by deprotection. To prepare 5,6-dehydro-PG₂s containing a hydrocarbon chain of the formula:

a monoalkyl copper complex or a dialkyl copper complex of the formula:

is subjected to 1,4-addition with 4R-t-butyldimethylsilyloxy-2-cyclopenten-1-one, and the resulting copper enolate is selzed with 6-carboalkoxy-1-iodo-2-hexyne or a derivative thereof.

[0052] PGs containing a methyl group instead of a hydroxy group at the C-11 position may be obtained as follows: PGA obtained by Jones oxidation of the hydroxy group at the C-9 position of the 11-osystate is allowed to react with a dimethyl copper complex to give II-dehydroxy.-Ie-methy-PGE. Attendively, an alcohol obtained after elimination of phenylbenzoyl group is converted to a tosylate. An unsaturated lactone obtained by DBU treatment of the tosylate is convented to a lactor. After introduction of an orachain using Wittig reaction, the resulting alcohol (C-9 position) is oxidized to give PGA. PGA is allowed to react with dimethyl copper complex to give 11-dehylbroxy. The resultinal

is reduced using sodium borohydride and the like to give II-dehydroxy-11-methyl-PGF.

[0053] PGs containing a hydroxymethly group instead of a hydroxyl group at the C-11 position is obtained as follow: 11-dehydroxy-11-hydroxymethly-FGE is obtained by a benzophenone-sensitized photoaddition of methanol to PGA. The resultant is, for example, reduced using sodium borohydride to give 11-dehydroxy-1-hydroxymethy-FGF.

[0054] 16-Fluoro-PGs may be obtained using dimethyl (3-fluoro-2-oxoalkyl)phosphonate anion in the preparation of an o.β-unsaturated ketone. Similarly, 19-methyl-PGs may be obtained using a dimethyl (6-methyl-2-oxoalkyl)phosphonate anion.

[0055] The preparations in the present invention are not construed to be limited to them, and suitable means for protection, oxidation, reduction and the like may be employed.

[0056] Examples of the preparation of the prostanoic acid compounds are described in the Japanese Patent Publications (unexamined) No. A-151552/1989, A-108/1990, A-96528/1990 and A-96529/1990.

[0057] The β-addresergic blockers and the prestancia call compounds used in the present invention can be used for the treatment of various disease and conditions of humans and animals in which lowering of ocular pressure is destrous and are usually administered systemically or topically by, for example, ophthalmic, oral, intravenous, subcutaneous, sectal administration set.

[0058] As used herein, the term "treatment" or "treating" refers to any means of control of a disease in a mammal, including preventing the disease, curing the disease, relieving the disease and arresting or relieving the development of the disease.

of the disease.

While the desage varies depending on the kind, age, weight, condition of the patient, such as humans or animals, severity of the disease, purpose of the treatment, judgement of the physician and route or period of administration, usually a satisfactory effect is obtained within the range of 0.01-500 µg/eye of the β-adrenergic blocker and 0.001-500 mg/kg of the prestancie acid compound.

[0060] The agents used in the present invention can be administered in the form of a pharmaceutical composition containing the active components and optionally other ingredients, such as carrier, diluent or excipient.

25 [0061] Such composition includes liquids such as ophthalmic solution, emulsion, dispersion etc. and semisolids such as gel, ointment etc.

(0062) Diluents for the aqueous solution or suspension include, for example, destilled water and physiological saline. Diluents for the nonaqueous solution and suspension include, for example, vegetable oils a g. cilie oil, [suitip paragine, mineral oil, and propylene glycol and p-octyldodecanol. The composition may also contain isotonization agents such as bodien bridder, doedline interest, codium critate, etc. to make isotonic with the lacinitian fluid and buffering agents such as borate buffer, phosphate buffer, etc. to maintain pH about 5.0 to 8.0. Further, stabilizers such as sodium sulfite, propylene glycol, etc., chelating agents such as sodium addated, etc., hickeners such as glycerot, carboxymethyled-luiose, carboxyminy loyhmer, etc. and preservatives such as methyl paraben, propyl paraben, etc may also be added, these can be sterilized etc. phy passing through a bacterial filter or ty heating.

35 [0063] The ophthalmic ointment may contain vaseline, Plastibase, Macropol, etc. as a base and surfactant for increasing hydrophilicity. It may also contain gelling agents such as carboxymethylcellulose, methylcellulose, carboxy-vinyl polymer, etc.

[0064] In addition, the composition may contain antibiotics such as chloramphenicol, penicillin, etc. in order to prevent or treat bacterial infection.

40 [0065] These composition may be packaged with an indication for administration. Such indication may be printing on package box, a bottle, a label, a separate paper sheet etc.

[0066] A more complete understanding of the present invention can be obtained by reference to the following Preparation Examples, Formulation Examples and Test Examples which are provided herein for purpose of illustration only and are not intended to limit the scope of the invention.

Preparations

[0067] Preparations of 13,14-dihydro-15-keto-20-ethyl-PGR₂ isopropyl ester, 13,14-dihydro-15-keto-20-ethyl-PGF₂ isopropyl ester and 13,14-dihydro-15-keto-20-ethyl-PGF₂α isopropyl ester (cf. Preparation chart I):

1) Preparation of 1S-2-oxa-3-oxo-6R-(3-oxo-1-trans-decenyl)-7R-(4-phenylbenzoyloxy)-cis-bicyclo[3.3.0]-octane

(3):

Commercially available (-)-Corey lactone (1) (7 g) was subjected to Collins oxidation in dichloromethane to

give aldehyde (2). The resultant was allowed to react with dimethyl (2-oxononyl)phosphonate (4.97 g) anion to give 15.2-oxa-3-oxo-6Ri-(3,3-ethylendloxy-1-trans-decenyl)-7R-(4-phenylbenzoyloxy)-cis-bicyclo(3.0)-octane (3).

 Preparation of 15-2-oxa-3-oxo-6R-(3-oxodecyl)-7R-(4-phenylbenzoyloxy)-cis-bicyclo[3.3.03-octane (4): Unsaturated ketone (3) (7.80 g) was reduced in ethyl acetate (170 ml) using 5% Pd/C under hydrogen atmos-

phere. The product obtained after the usual work-up (4) was used in the following reaction.

3) Preparation of 1S-2-oxa-3-oxo-6R-(3,3-ethylenedioxy-decyl)-7R-(4-phenylbenzoyloxy)-cis-bicyclo[3,3,0]-octane (5):

Saturated ketone (4) was converted to ketal (5) in dry benzene (150 ml) using ethylene glycol and p-toluenesulfonic acid (catalytic amount).

4) Preparation of 1S-2-oxa-3-oxo-5R-(3,3-ethylenedioxy-decyl)-7R-hydroxy-cis-blcyclo[3,3,0]-octane (6):

To a solution of ketal (5) in absolute methanol (150 ml) was added poissium carbonate (2.73 g). The mixture was attired overight at from temperature. After neutralization with acetic acid, the resultant was concentrated under reduced pressure. The resulting trute product was entrated with eithy acetale. The organic layer was washed with a dilute aqueous solution of sodium biotochonate and a saline, and dried. The crude product obtained she revolved to the product obtained she revolved to the contract product of the contract product product of the contract product produc

Alcohol (6) (0.80 g) was reduced in dry toluena (8 ml) using DIBAL-H at -78 °C to give lectol (7).

Preparation of 13,14-dhydro-15,15-ethylenedioxy-20-ethyl-PGF₂α (8):

A DMSO solution of lactol (7) was added to ylide prepared from (4-carboxybuty)triphenylphosphonium bromide (3.65 g). The reaction mixture was silred overnight to give carboxylic acid (8). 7) Preparation of 13,14-dinydro

Carboxylic acid (8) was converted to 13,14-dihydro-15,15-ethylenedioxy-20-ethyl-PGF₂α isopropyl ester (9) using DBU and isopropyl iodide in acetonitrile.

Yield; 0.71 g

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Preparation of 13,14-dihydro-15-keto-20-ethyl-PGF₂α isopropyl ester (10):

13.14-dhydro-15,15-ethylenedoxy-20-ethyl-PGF₂x isopropyl ester (9) (0.71 g) was kept in acetic acid/THF/ water (3/17) at 40 °C for 3 hours. The crude product obtained after concentration under reducted pressure was chromatographed to give 13,14-dihydro-15-keto-20-ethyl-PGF₂x isopropyl ester (10). Yelet 0.554 a

Preparation of 13,14-dihydro-15-keto-20-ethyl-PGA₂α isopropyl ester (12):

A solution of 13,14-dihydro-15-keto-20-ethyl-PGF₂ct isopropyl ester (10) (0.125 g) and p-tolucnesulfonyl chloride (0.112 g) in pyridine (5 ml) was maintained at 0 °C for 2 days. According to the usual work-up, tosylate (11) was obtained.

Tosylate (11) was subjected to Jones oxidation in acetone (8 ml) at -25 °C. The crude product obtained after the usual work-up was chromatographed to give 13,14-dihydro-15-keto-20-ethyl-PGA₂α isopropyl ester (2). Yeld: 0.050 α

Preparation of 13,14-dihydro-15,15-ethylenedioxy-20-ethyl-11-t-butyldimethylsiloxy-PGF₂α isopropyl ester

dimethylomemide (25 ml), butyldimethyteliy chloride (1089 g) and midazole (0.49 g) was dissolved in dry N,Ndimethylomemide (25 ml), butyldimethyteliy chloride (1089 g) and midazole (0.49 g) was added thereto. The resultant was stirred at room temperature overnight. The resction mixture was concentrated under reduced. The sure, and the resulting crude product was chromatographed to give 13,14-dihydro-15,15-ethylenedioxy-20-ethyl-11-butyldimethylsiony-PGF-ça ilcoprovel setal (13).

Yield; 2.641 g

11) Preparation of 13,14-dihydro-15,15-ethylenedioxy-20-ethyl-11-t-butyldimethylsiloxy-PGE₂ isopropyl ester (14): 13,14-dihydro-15,15-ethylenedioxy-20-ethyl-11-butyldimethylsiloxy-PGE₂ is logropyl ester (13) (1.257 g) was subjected to Jones oxidation at 40°C. After the usual work-up, the resulting crude product was chromatographed to give 13,14-dihydro-15,15-ethylenedioxy-20-ethyl-11-butyldimethylsiloxy-PGE₂ isopropyl ester (14). Yiled; 1.082 g)

12) Preparation of 13,14-dihydro-15-keto-20-ethyl-PGE2 isopropyl ester (15):

To a solution of 13,14-dihydro-15,15-ethylene-dioxy-20-ethyl-11-butyldimethylsiloxy-PGE₂α isopropyl ester (14) in acetonitrile was added hydrolluoric acid (46% aqueous solution). The mixture was stirred at room temperature for 40 minutes. The condeproducts obtained after usual work-up was chromatographed to give 13,14-dihydro-15-keto-20-ethyl-PGE₂ isopropyl ester (15).

Yield; 0.063 g (97%)

Formulation Example 1	
Timolol maleate	0.1 g
Physiological saline	q s. to 100 ml

Formulation Example 2	
13.14-dihydro-15-keto-20-ethyl-PGF ₂ α isopropyl ester	0.01 g
Nonion Surfactant	1.0 g
Physiological saline	q.s. to 100 ml

Test Example 1

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[066] Hypotensive effect of Timolol was evaluated in the enhancement phase of aqueous humour production and the suppression phase of aqueous humour production of rabbits. Since the circadian rhythm of rabbits, different from that of humans, has the enhancement phase of aqueous humour production at night and the suppression phase of aqueous humour production at deytime, the following two experiments were performed.

(1) Enhancement phase of aqueous humour production:

[0069] White rabbits (n=8) were used in the experiment of intraocular pressure measurement after keeping under the environmental conditions including a light and darkness cycle consisting of a light period from 21:00 to 9:00 and dark period from 9:00 to 21:00 for more than one week. In the experiment, 55 µ d at 0.5% "implole yetrop (Trademark: Timoptol) was administered to one eye at 11:00 (dark time). The coular tension was measured immediately before and 1 hour after the administration and the difference between the obtained two values was expressed as decrease in intraocular pressure (alOP).

30 (2) Suppression phase of aqueous humour production:

[0070] White rabbits (n=12) were used in the experiment of intracoular pressure measurement after keeping under the environmental conditions including a light and darkness cycle consisting of a light period from 8 00 to 8 00 for more than one week. In the experiment, 55 µd of a 0.5% Timolol eyed op (Trademark: Timopto) was administered to one eye at 10:00 (light time). The coular tension was measured immediately before and 3 hours after the administration and the difference between the obtained two values was expressed as decrease in intracoular pressure (aIOP). The results are shown in Table 1.

| Table 1 | | Enhancement Phase* | Suppression Phase* | AIOP (mmHg) | 6.4±1.0 | 2.5±0.8 |

· Production of aqueous humou

g [0071] Then, the procedure of the experiment (2) was repeated except that a 0.12% eye drop of 13,14-dihydro-15-keto-20-ethyl-PGF₂α isopropyl ester was used in place of the 0.5% Timobil eye drop. The results are shown in Table 2.

	Table 2
	Suppression Phases
ΔIOP (mmHg)	7.1±0.7

See footnole of Table

55 Tesl Example 2

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[0072] A 0.5% Timolol eye drop was intraocularly administered to subjects of glaucoma (n=8) twice (morning and evening) a day for 4 weeks. Differences in intraocular pressure were measured as in Test Example 1 and expressed

as decrease in intraocular pressure (AIOP). The results are shown in Table 3.

Table 3

	Enhancement Phase* (11:00)	Suppression Phase+ (19:00)
ΔIOP (mmHg)	2.9±0.8	0.4±0.7

See footnote of Table 1.

[0073] Separalely, the above experiment was repeated using subjects of glaucoma (n=10) and administering a 0.12% eye drop of 13,14-dihydro-15-keto-20-ethyl-PGF₂α isopropyl ester in place of the 0.5% Timolol eye drop and decrease in intraocular pressure (a10P) was determined at the suppression phase of a queous humour production (19:00). The results are shown in Table 4.

	Suppression Phase*
ΔIOP (mmHg)	2.1±0.3

See footnote of Table 1.

Claims

- The use of a β-adrenergic blocker and of a derivative of prostanoic acid for the manufacture of a therapeutic kit for the concomitant treatment of ocular hypertension, wherein the β-adrenergic blocker is to be administered only in the enhancement phase of aqueous humour production and the derivative of prostanoicacid is to be administered only in the suppression phase of aqueous humour production.
- 2. The use according to claim 1 wherein the derivative of prostanoic acid is represented by the following formula (I): -

$$\begin{array}{c}
X \\
R_1 \longrightarrow A \\
B \longrightarrow C \longrightarrow R_2
\end{array}$$
(1)

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X and Y are hydrogen, hydroxy, halo, lower allyl, hydroxy(lower)alkyl or oxo, with the proviso that at least one of X and Y is a group other than hydrogen, and 5-membered ring may have at least one double bond, A is -COOH or its pharmaceutically sall or ceter, B is -Chi₂-Chi₂--Chi-Chi-C o-CarC.

Z is

wherein R3 is lower alkyl or acyl,

 R_1 is bivalent saturated or unsaturated, lower or medium aliphatic hydrocarbon residue which is unsubstituted or substituted with halo, oxo or anyl,

R₂ is saturated or unsaturated, medium aliphatic hydrocarbon residue which is unsubstituted or substituted with halo, hydroxy, oxo, lower alkoxy, lower alkanoyloxy, cyclo(lower)alkyl, aryl or aryloxy.

3. The use according to claim 1 wherein the derivative of prostanoic acid is represented by the following formula (II): -

$$R_1-A$$
 $B-CO-R_2$
(II)

wherein

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- X and Y are hydrogen, hydroxy, halo, lower allkyl, hydroxy(lower)alkyl or oxo, with the proviso that at least one of X and Y is a group other than hydrogen, and 5-membered ring may have at least one double bond, A is COOH or its pharmaceutically acceptable sall or estel, is -CH₂-CH₂.
- -CH=CH- or -C=C, R_i is bivalent saturated or unsaturated, lower or medium aliphatic hydrocarbon residue which is unsubstituted or substituted with halo, oxo or aryl, R_i is saturated or unsaturated, medium aliphatic hydrocarbon residue having 5 or more carbon atoms in the main or straight chain molety which is unsubstituted or substituted with halo, hydroxy, oxo, lower alikoxy, lower alikanoyloxy, cyclotlower/alikyl, aryl or aryloxy.
- The use according to claim 3 wherein the derivative of prostanoic acid is 13,14-dihydro-15-keto-20-ethyl-PGF_{2a} isopropyl ester.
 - 5. The use according to any one of claims 1 to 3 wherein the derivative of prostanoic acid is a prostaglandin.
 - 6. The use according to claim 5 wherein the derivative of prostanoic acid is a prostaglandin F.
 - 7. The use according to claim 5 wherein the derivative of prostanoic acid is a 15-keto-prostaglandin.
 - The use according to any one of claims 1 to 7 wherein the β-adrenergic blocker is selected from the group consisting
 of Timolol, Befundol, Betaxolol, Levobundol, Carteolol and pharmaceutically acceptable salts thereof.
 - The use of a β-adrenergic blocker and of a derivative of prostancic acid for the manufacture of a therapeutic kit
 for the concomitant treatment of glaucoma, wherein the β-adrenergic blocker is to be administered only in the
 enhancement phase of aqueous humour production and the derivative of prostancic acid is to be administered
 only in the suppression phase of aqueous humour production.
 - 10. The use according to claim 9 wherein the derivative of prostanoic acid is represented by the following formula (I):-

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$$\begin{array}{c}
X \\
R_1 - A \\
B - C - R_2
\end{array}$$
(J)

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X and Y are hydrogen, hydroxy, halo, lower alkyl, hydroxy(lower)alkyl or oxo, with the proviso that at least one of X and Y is a group other than hydrogen, and 5-membered ring may have at least one double bond, A is -COOH or its pharmaceutically acceptable salt or ester, B is -CH2-CH2-,-CH=CH- or -C=C-, Z is

wherein R3 is lower alkyl or acyl,

R₁ is bivalent saturated or unsaturated, lower or medium aliphatic hydrocarbon residue which is unsubstituted or substituted with halo, oxo or aryl,

R2 is saturated or unsaturated, medium aliphatic hydrocarbon residue which is unsubstituted or substituted with halo, hydroxy, oxo, lower alkoxy, lower alkanoyloxy, cyclo(lower)alkyl, aryl or aryloxy.

11. The use according to claim 9 wherein the derivative of prostanoic acid is represented by the following formula (II):

$$R_1$$
—A (II)

X and Y are hydrogen, hydroxy, halo, lower alkyl, hydroxy(lower)alkyl or oxo, with the proviso that at least one of X and Y is a group other than hydrogen, and 5-membered ring may have at least one double bond, A is - COOH or its pharmaceutically acceptable salt or ester, B is -CH2-CH2-, -CH=CH- or -C=C-, R1 is bivalent saturated or unsaturated, lower or medium aliphatic hydrocarbon residue which is unsubstituted or substituted with halo, oxo

or anyl, R₂ is saturated or unsaturated, medium aliphatic hydrocarbon residue having 5 or more carbon atoms in the main or straight chain molely which is unsubstituted or substituted with halo, hydroxy, oxo, lower alixoxy, lower alkanoyloxy, oxo(clower) alixyl, anyl or anyloxy.

- The use according to claim 11 wherein the derivative of prostanoic acid is 13,14-dihydro-15-keto-20-ethyl-PGF_{2α} isopropyl ester.
 - 13. The use according to any one of claims 9-11 wherein the derivative of prostanoic acid is a prostaglandin.
- 14. The use according to claim 13 wherein the derivative of prostanoic acid is a prostaglandin F.
 - 15. The use according to claim 13 wherein the derivative of prostanoic acid is a 15-keto-prostaglandin.
- 16. The use according to any one of claims 9 to 15 wherein the β-adrenergic blocker is selected from the group consisting of Timolol, Betunolol, Betaxolol, Levobunolol, Carteolol and pharmaceutically acceptable salts thereof.
 - 17. The use according to any preceding claim wherein the enhancement phase is in the daytime and the suppression phase is at night.

Patentansprüche

- Verwendung eines
 ß-adrenergen Blockers und eines Derivats von Prostansäure zur Herstellung eines therapeutschen Klis für die gleichzeitige bzw. begleitende Behandlung von Augenhochdruck, wobei der
 ß-adrenerge Blokern und in der Verstärkungsphase der Kammerwasserproduktion verabreicht werden soll und das Derivat der Prostensäugen nur in der Unterdückungsphase der Kammerwasserproduktion verabreicht werden soll.
 - Verwendung nach Anspruch 1, dadurch gekennzeichnet, dass das Derivat der Prostansäure durch die folgende allgemeine Formel (I) angegeben wird:

worin

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X und Y für Wasserstoff, Hydroxy, Halogen, Niedrigaßkyl, Hydroxy(niedrig)alkyl oder Oxo stehen, mit der Magheb, dass mindestens eines von X und Y eine andere Gruppe als Wasserstoff ist, wobei der 5-gliedrige Ring mindestens eine Doppebindung haber kann, A für COOH oder das pharmazeutischen Satz oder den pharmazeutischen Ester davon steht, B für -CH₂-CH₂-C-H=CH oder -C=C steht, Zfür

steht.

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wobai R₃ die Bedeutung Niedrigalkyl oder Acyl hat,

R₁ für einen zweiwertigen gesättigten oder ungesättigten, niederen oder mittleren allphatischen Kohlenwasserstoffrest, der unsubstituiert ist oder mit Halogen, Oxo oder Aryl substituiert ist, steht,

R₂ für einen gesättigen oder ungesättigten mittleren aliphatischen Kohlenwasserstoffrest, der unsubstituiert ist oder mit Halogen, Hydroxy, Oxo, Niedrigalkoxy, Niedrigalkanoyloxy, Cyclo(niedrig)alkyl, Aryl oder Aryloxy substituiert ist, steht.

 Verwendung nach Anspruch 1, dadurch gekennzeichnet, dass das Derivat der Prostansäure durch die tolgende Formel (III):

$$R_1$$
—A (II)

angegeben wird, worin

X und Y für Wassenstoff, Hydroxy, Helogen, Niedrigsleyh, Hydroxy/Iniedrigsleyh oder Oxo stehen, mit der Magglabe, dass mindestens eines von X und Y eine andere Gruppe als Wassensroll sit, wobei der z-sjelerdige Ring mindestens eine Doppebindung haben kann, A für -CDOH oder das pharmarsutsleche Salz oder den pharmarsutslechen salz oder den pharmarsutslechen zu zu der den zu zu zu zugestätigten oder ungestätigten oder mitteren allphatischen Kohlenwasserstoffrest, der unsubstitutient ist oder im Halogen, Oxo oder Ard substitutier ist, sicht in

R, für einen gesätigien oder ungesättigen, mittleren aliphatischen Kohlenwasserstoffrest mit 5 oder mehr Kohlenstoffatomen in der Hauptkeite oder der geradektigen Grupplerung, der unsubstituler ist oder mit Habgen, Hydroxy, Cxx, Niedrigalikoxy, Niedrigalikanopixox, Cyteloniedrigalikyl, Aryl oder Anjors substituler ist, sieht.

- Verwendung nach Anspruch 3, dadurch gekennzeichnet, dass das Derivat der Prostansäure der 13,14-Dihydro-15-keto-20-ethyl-PGF_{2u}-isopropylester ist.
- Verwendung nach einem der Ansprüche 1 bis 3, dadurch gekennzeichnet, dass das Derivat der Prostansäure Prostaglandin ist.
 - 6. Verwendung nach Anspruch 5, dadurch gekennzeichnet, dass das Derivat der Prostansäure Prostaglandin Fist.
- Verwendung nach Anspruch 5, dadurch gekennzeichnet, dass das Derivat der Prostansäure ein 15-Ketoprostaglandin ist.
- Verwendung nach einem der Ansprüche 1 bis 7, dadurch gekennzeichnet, dass der β-adrenerge Blocker aus der Gruppe, bestehend aus Timolol, Belavolol, Belavolol, Levobunolol, Carleolol und den pharmazeutisch annehmberen Satzen devon ausgewählt wird.
- Verwendung eines β-adrenergen Blockers und eines Derivats der Prostansäure zur Herstellung eines therapeutschen Kils für die gleichz eilige bzw. begiebende Behandung des Gleukoms, wobei der β-adrenerge Blocker nur in der Verstätkungsphase der Kammerwasserproduktion verderbeitwirteden soll und das Derivat der Prostansäure

nur in der Unterdrückungsphase der Kammerwasserproduktion verabreicht werden soll.

 Verwendung nach Anspruch 9, dadurch gekennzeichnet, dass das Derivat der Prostansäure durch die folgende allgemeine Formel (f) angegeben wird:

$$\begin{array}{c}
X \\
R_1 \longrightarrow A
\end{array}$$

$$\begin{array}{c}
R_1 \longrightarrow A \\
B \longrightarrow C \longrightarrow R_2 \\
Z
\end{array}$$

$$\begin{array}{c}
X \\
Z
\end{array}$$

worin

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X und Y für Wasserstoff, Hydroxy, Halogen, Niedrigalkyl, Hydroxy(niedrigalkyl oder Oxo stehen, mit der Menden Gruppe als Wasserstoff st, wobel der 5-gleedrige Ring mindestens eine Doppebindung haber kann, A für COOD ded rad sey harmzeut/siche Salz oder den pharmazeu-tischen Ester davon steht, B für -CH₂-CH₂-C-H=CH oder -C=C steht, Z für

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- steht, wobei R₃ die Bedeutung Niedrigalkyl oder Acyl hat,
- R₁ für einen zweiwertigen gesättigten oder ungesättigten, niederen oder mittleren aliphatischen Kohlenwasserstoffrest, der unsubstituiert ist oder mit Halogen, Oxo oder Aryl substituiert ist, steht,
- R₂ für einen gesättigen oder ungesättigten mittleren allphatischen Kchlenwasserstoffrest, der unsubslitulert iser mit Halogen, Hydroxy, Oxo, Niedrigalkoxy, Niedrigalkanoyloxy, Cyclo(niedrigalky), Aryl oder Aryloxy substituiert ist, steht.
- Verwendung nach Anspruch 9, dadurch gekennzeichnet, dass das Derivat der Prostansäure durch die folgende Formel (II):

angegeben wird,

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worin

X und Y für Wassenstoff, Hydroxy, Halogen, Niedrigalbyl, Hydroxy/Ineidrigalbity, oder Oxo stehen, mit der Maßgabe, disse miniestens eines von X und Y eine andere Gruppe auf Wissenstoff ist, wobei der 7-spliedrige Ring mindestens eine Doppelbindung haben kann. A für -COOH oder des pharmazeutsiche Salz oder den pharmazeutsichen Salz oder son steht, B für -CHO-QH-, C-Me-CHO-QH- oder -C-P cehr, R, für ehen zweitwertigen gesättigen oder ungesättigten niederen oder mitteren allphatischen Kohlenwassersbürfers, der unsubstütigen ist oder mit Halogen, Oxo oder Ard substütiger ist, selbt.

R, für einen gesättigten oder ungesättigen, mittleren allphatischen Kohlenwasserstoffrest mit 5 oder mehr Kohlenstoffatomen in der Haupkkette oder der geranketitigen Gruppierung, der unsubstituler it at oder mit Hatogen, Hydroxy, Oxo, Niedrigalkoxy, Niedrigalkanoyloxy, Cyclofinedrigalkyi, Ayl oder Anjorky substituler it sit, sehb.

- Verwendung nach Anspruch 11, dadurch gekennzeichnet, dass das Derivat der Prostansäure 13,14-Dihydro-15-keto-20-ethyl-PGF_{2n}-isopropylester ist.
 - Verwendung nach einem der Ansprüche 9 bis 11, dadurch gekennzeichnet, dass das Derivat der Prostansäure Prostaglandin ist.
- 14. Verwendung nach Anspruch 13, dadurch gekennzelchnet, dass das Derivat der Prostansäure ein Prostaglandin F ist.
 - Verwendung nach Anspruch 13, dadurch gekennzeichnet, dass das Derivat der Prostansäure 15-Ketoprostaglandin ist.
 - 16. Verwendung nach einem der Ansprüche 9 bis 15, dadurch gekennzeichnet, dass der β-adrenerge Blocker aus der Gruppe, bestehend aus Timolol, Beltaxolol, Levobunolol, Carteolol und den pharmazeutisch annehmberen Salzen davon, ausgewählt wird.
- Verwendung nach einem der vorstehenden Ansprüche, dadurch gekennzeichnet, dass die Verstärkungsphase tagsüber erfolgt und dass die Unterdrückungsphase über Nacht erfolgt.

Revendications

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- 1. Utilisation d'un inhibiteur β-adrénergique et d'un dérivé d'acide prostanoique pour la fabrication d'une trousse thérapeutique destinée au traitement concomitant de thypertension oculaire, dans laquelle l'inhibiteur β-adrénergique n'est à definisiter que dans la phase de stimulation de la production d'humeur aqueuse et le détrivé d'acide prostanoique n'est à administrer que dans la phase de frénàge de la production d'humeur aqueuse.
- Utilisation selon la revendication 1 dans laquelle le dérivé d'acide prostanoîque est représenté par la formule (i) suivante:

dans laquelle

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X et Y sort des atomes d'hydrogène ou des groupes hydroxy, halogéne, altyle inférieur, hydroxyaltyle inférieur ou oxo, à condition qu'au moins un des radicaux X et Y soit un groupe autre qu'un atome d'hydrogène, et le noyeu à 5 chaitons peut comporter au moins une double liaison. A est -COOH ou son sel ou ester pharmaceutiquement acceptable, B est -CH₂-CH₂-, CH=CH- ou -C=C-. Zest

où R₃ est un groupe alkyle inférieur ou acyle,

R₁ est un résidu hydrocarboné allphatique bivalent, inférieur ou moyen, saturé ou insaturé, qui est non substitué ou substitué par un groupe halogéno, oxo ou aryle,

R₂ est un résidu hydrocarboné aliphatique moyen, saturé ou insaturé, qui est non substitué ou substitué par un groupe halogéno, hydroxy, oxo, alcoxy inférieur, alcanoyloxy, cycloalkyle inférieur, aryle ou aryloxy.

3. Utilisation selon la revendication 1 dans laquelle le dérivé d'acide prostanoïque est représenté par la formule (II) suivante:

$$R_1$$
—A (II)

dans laquelle

X et Y sont des atomes d'hydrogène ou des groupes hydroxy, halogéno, alkyle inférieur, hydroxyalkyle inférieur ou oxo, à condition qu'au moins un des radicaux X et Y soit un groupe autre qu'un atome d'hydrogène, et le noyeu à 5 chaînons peut comporter au moins une double liaison. A est -CODH ou son set ou este pharmaceutiquement acceptable, B est -CH₂-CH₂-CH-CH-CH- ou -C=C-C. R₁ est un résidu hydrocarboné aliphatique bivalent, inférieur ou moyen, saturé ou insaturé, qui est non substitué ou substitué par un groupe halogéno, oxo ou aryle, R₂ est un résidu hydrocarboné aliphatique moyen, saturé ou insaturé, comportant au moins 5 atomes de carbone dans le groupement principal ou à chaîne inleaire, qui est non substitué ou substitué par un groupe halogéno, hydroxy, oxo, altoxy inférieur, actenovioys inférieur, cyclosikyle inférieur, arvice ou aryloxy.

- Utilisation seion la revendication 3 dans laquelle le dérivé d'acide prostancique est l'ester isopropylique de 13,14-dihydro-15-céto-20-éthyl-PGF_{2u}.
- Utilisation selon l'une quelconque des revendications 1 à 3 dans laquelle le dérivé d'acide prostanoïque est une prostagiandine.
 - 6. Utilisation selon la revendication 5 dans laquelle le dérvé d'acide prostanoïque est une prostaglandine F.
- Utilisation selon la revendication 5 dans laquelle le dérivé d'acide prostanoïque est une 15-céto-prostaglandine.
 - Utilisation selon l'une quelconque des revendications 1 à 7 dans laquelle l'inhibiteur β-adrénergique est choisi dans le groupe constitué par le Timolol, le Béfunolol, le Béfunolol, le Lévobunoloi, le Cartéolol et leurs sels pharmaceutiquement acceptables.
- 9 Utilisation σ'un inhibiteur β-adrénergique et d'un dérivé d'acide prostanoique pour la fabrication d'une trousse thérapeutique destinée au traitement concomitant du glaucome, dans laquelle l'inhibiteur β-adrénergique n'est à administrer que dans la phase de stimulation de la production d'humeur aqueuse et le dérivé d'acide prostanoïque n'est à administrer que dans la phase de le freinage de la production d'humeur apueuse.
- 25 10. Utilisation selon la revendication 9 dans laquelle le dérivé d'acide prostanoïque est représenté par la formule (I) suivante:

$$\begin{array}{c}
X \\
R_1 \longrightarrow A
\end{array}$$

$$\begin{array}{c}
R_1 \longrightarrow A \\
\end{array}$$

$$\begin{array}{c}
C \longrightarrow R_2 \\
Z
\end{array}$$

$$\begin{array}{c}
Z \longrightarrow R_2
\end{array}$$

45 dans laquelle

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X et Y sont des atomes d'hydrogène ou des groupes hydroxy, halogène, alkyle intérieur, hydroxyalkyle intérieur ou oxo, à condition qu'au moins un des radicaux X et Y soit un groupe autre qu'un atome d'hydrogène, et le noyau às Chainions pout comporter au moins une double lisison. A est -COOH ou son sel ou ester pharmaceutiquement acceptable, B est -CH₂-CH₂-, -CH=CH- ou -C=C-, Z est

où R₃ est un groupe alkyle inférieur ou acyle,

R₁ est un résidu hydrocarboné aliphatique bivalent, inférieur ou moyen, saturé ou insaturé, qui est non substitué ou substitué par un groupe halogéno, oxo ou aryle,

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R₂ est un résidu hydrocarboné aliphatique moyen, saturé ou insaturé, qui est non substitué ou substitué par un groupe halogèno, hydroxy, oxo, alcoxy inférieur, alcanoyloxy inférieur, cycloalkyle inférieur, aryle ou aryloxy.

11. Utilisation selon la revendication 9 dans laquelle le dérivé d'acide prostanoïque est représenté par la formule (II) suivante:

$$R_1$$
—A (II)

25 dans laquelle

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- uans lacquere X et Y sont des atomes d'hydrogène ou des groupes hydroxy, halogéno, alkyle inlérieur, hydroxyalkyle intérieur ou cox, à condition qu'au moins un des radicaux X et Y soit un groupe autre qu'un atome d'hydrogène, et le noyau à 5 chaînons peut comporter au moins une double liaison, A est COOH ou son set ou estre pharmaceutiquement acceptable, B est CHy-CHy, -CH-CH- ou -CH-C, R, est un résidu hydrocertoné aliphatique bivalent, intérieur ou moyen, saturé ou insaturé, qui est non substitué ou substitué par un groupe halogène, cox ou a nyle, R, est un résidu hydrocarboné aliphatique moyen, saturé ou insaturé, comportant au moins 5 atomes de carbone dans le groupement principal ou à chaîne linéaire, qui est non substitué ou substitué par un groupe halogène, hydroxy, oxe, aboxy inférieur, ateanolyony inférieur, cyclosihyle inférieur, argie ou sylroxy.
- Utilisation selon la revendication 11 dans laquelle le dérivé d'acide prostanoïque est l'ester isopropylique de 13,14-dihydro-15-céto-20-éthyl-PGF_{2a}.
 - 13. Utilisation selon l'une quelconque des revendications 9 à 11 dans laquelle le dérivé d'acide prostanoïque est une prostaglandine.
 - 14. Utilisation selon la revendication 13 dans laquelle le dérivé d'acide prostanoïque est une prostaglandine F.
 - 15. Utilisation selon la revendication 13 dans laquelle le dérivé d'acide prostanoïque est une 15-céto-prostaglandine.
- 5 16. Utilisation seton l'une quelconque des revendications 9 à 15 dans laquelle l'inhibiteur β-adrénergique est choisi dans le groupe constitué par le Timolol, le Béfunolol, le Bétaxolol, le Lévobunolol, le Cartéolol et leurs sels pharmaceutiquement accerdables.
- 17. Utilisation seton l'une quelconque des revendications précédentes, dans laquelle la phase de stimulation est dans la journée et la phase de freinage est la nuit.

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